# REMOVAL OF OXYGEN FROM METAL OXIDES AND SOLID SOLUTIONS BY ELECTROLYSIS IN A FUSED SALT

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#### Abstract of WO9964638

A method for removing a substance (X) from a solid metal or semi-metal compound (M<1>X) by electrolysis in a melt of M<2>Y, comprises conducting the electrolysis under conditions such that reaction of X rather than M<2> deposition occurs at an electrode surface, and that X dissolves in the electrolyte M<2>Y. The substance X is either removed from the surface (i.c. M<1>X) or by means of diffusion extracted from the care material. The temperature of the fused salt is chosen below the melting temperature of the metal M<1>. The potential is chosen below the decomposition potential of the electrolyte.

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#### (57) Abstract

A method for removing a substance (X) from a solid metal or semi-metal compound ( $M^{I}X$ ) by electrolysis in a melt of  $M^{2}Y$ , comprises conducting the electrolysis under conditions such that reaction of X rather than  $M^{2}$  deposition occurs at an electrode surface, and that X dissolves in the electrolyte  $M^{2}Y$ . The substance X is either removed from the surface (i.c.  $M^{I}X$ ) or by means of diffusion extracted from the care material. The temperature of the fused salt is chosen below the melting temperature of the metal  $M^{1}$ . The potential is chosen below the decomposition potential of the electrolyte.

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# REMOVAL OF OXYGEN FROM METAL OXIDES AND SOLID SOLUTIONS BY ELECTROLYSIS IN A FUSED SALT

#### Field of Invention

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This invention relates to a method for reducing the level of dissolved oxygen or other elements from solid metals, metal compounds and semi-metal compounds and alloys. In addition, the method relates to the direct production of metal from metal oxides or other compounds.

### Background to the Invention

Many metals and semi-metals form oxides, and some have a significant solubility for oxygen. In many cases, the oxygen is detrimental and therefore needs to be reduced or removed before the metal can be fully exploited for its mechanical or electrical properties. For example, titanium, zirconium and hafnium are highly reactive elements and, when exposed to oxygen-containing environments rapidly form an oxide layer, even at room temperature. This passivation is the basis of their outstanding corrosion resistance under oxidising conditions. However, this high reactivity has attendant disadvantages which have dominated the extraction and processing of these metals.

As well as oxidising at high temperatures in the conventional way to form an oxide scale, titanium and other elements have a significant solubility for oxygen and other metalloids (e.g. carbon and nitrogen) which results in a serious loss of ductility. This high reactivity of titanium and other Group IVA elements extends to reaction with refractory materials such as oxides, carbides etc. at elevated temperatures, again contaminating and embrittling the basis metal. This behaviour is extremely deleterious in the commercial extraction, melting and processing of the metals concerned.

Typically, extraction of a metal from the metal oxide is achieved by heating the oxide in the presence of a reducing agent (the reductant). The choice of reductant is determined by the comparative thermodynamics of the oxide and the reductant, specifically the free energy balance in the reducing reactions. This balance must be negative to provide the driving force for the reduction to proceed.

The reaction kinetics are influenced principally by the temperature of reduction and additionally by the chemical activities of the components involved. The latter is often an important feature in determining the efficiency of the process and the completeness of the reaction. For example, it is often found that although

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this reduction should in theory proceed to completion, the kinetics are considerably slowed down by the progressive lowering of the activities of the components involved. In the case of an oxide source material, this results in a residual content of oxygen (or another element that might be involved) which can be deleterious to the properties of the reduced metal, for example, in lower ductility, etc. This frequently leads to the need for further operations to refine the metal and remove the final residual impurities, to achieve high quality metal.

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Because the reactivity of Group IVA elements is high, and the deleterious effect of residual impurities serious, extraction of these elements is not normally carried out from the oxide, but following preliminary chlorination, by reducing the chloride. Magnesium or sodium are often used as the reductant. In this way, the deleterious effects of residual oxygen are avoided. This inevitably leads, however, to higher costs which make the final metal more expensive, which limits its application and value to a potential user.

Despite the use of this process, contamination with oxygen still occurs. During processing at high temperatures, for example, a hard layer of oxygen-enriched material is formed beneath the more conventional oxide scale. In titanium alloys this is often called the "alpha case", from the stabilising effect of oxygen on the alpha phase in alpha-beta alloys. If this layer is not removed, subsequent processing at room temperature can lead to the initiation of cracks in the hard and relatively brittle surface layer. These can then propagate into the body of the metal, beneath the alpha case. If the hard alpha case or cracked surface is not removed before further processing of the metal, or service of the product, there can be a serious reduction in performance, especially of the fatigue properties. Heat treatment in a reducing atmosphere is not available as a means of overcoming this problem because of the embrittlement of the Group IVA metals by hydrogen and because the oxide or "dissolved oxygen" cannot be reduced or minimised. The commercial costs of getting round this problem are significant.

In practice, for example, metal is often cleaned up after hot working by firstly removing the oxide scale by mechanical grinding, grit-blasting, or using a molten salt, followed by acid pickling, often in HNO<sub>3</sub>/HF mixtures to remove the oxygen-enriched layer of metal beneath the scale. These operations are costly in terms of loss of metal yield, consumables and not least in effluent treatment. To minimise scaling and the costs associated with the removal of the scale, hot

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working is carried out at as low a temperature as is practical. This, in itself, reduces plant productivity, as well as increasing the load on the plant due to the reduced workability of the material at lower temperatures. All of these factors increase the costs of processing.

In addition, acid pickling is not always easy to control, either in terms of hydrogen contamination of the metal, which leads to serious embrittlement problems, or in surface finish and dimensional control. The latter is especially important in the production of thin materials such as thin sheet, fine wire, etc.

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It is evident therefore, that a process which can remove the oxide layer from a metal and additionally the dissolved oxygen of the sub-surface alpha case, without the grinding and pickling described above, could have considerable technical and economic benefits on metal processing, including metal extraction.

Such a process may also have advantages in ancillary steps of the purification treatment, or processing. For instance, the scrap turnings produced either during the mechanical removal of the alpha case, or machining to finished size, are difficult to recycle due to their high oxygen content and hardness, and the consequent effect on the chemical composition and increase in hardness of the metal into which they are recycled. Even greater advantages might accrue if material which had been in service at elevated temperatures and had been oxidised or contaminated with oxygen could be rejuvenated by a simple treatment. For example, the life of an aero-engine compressor blade or disc made from titanium alloy is constrained, to a certain extent, by the depth of the alpha case layer and the dangers of surface crack initiation and propagation into the body of the disc, leading to premature failure. In this instance, acid pickling and surface grinding are not possible options since a loss of dimension could not be tolerated. A technique which lowered the dissolved oxygen content without affecting the overall dimensions, especially in complex shapes, such as blades or compressor discs, would have obvious and very important economic benefits. Because of the greater effect of temperature on thermodynamic efficiency these benefits would be compounded if they allowed the discs to operate not just for longer times at the same temperature, but also possibly at higher temperatures where greater fuel efficiency of the aeroengine can be achieved.

In addition to titanium, a further metal of commercial interest is Germanium, which is a semi-conducting metalloid element found in Group IVA of the Periodic

Table. It is used, in a highly purified state, in infra-red optics and electronics. Oxygen, phosphorus, arsenic, antimony and other metalloids are typical of the impurities which must be carefully controlled in Germanium to ensure an adequate performance. Silicon is a similar semiconductor and its electrical properties depend critically on its purity content. Controlled purity of the parent silicon or germanium is fundamentally important as a secure and reproducible basis, onto which the required electrical properties can be built up in computer chips, etc.

US Patent 5,211,775 discloses the use of calcium metal to deoxidise titanium. Okabe, Oishi and Ono (Met. Trans B. 23B (1992):583, have used a calcium-aluminium alloy to deoxidise titanium aluminide. Okabe, Nakamura, Oishi and Ono (Met. Trans B. 24B (1993):449) deoxidised titanium by electrochemically producing calcium from a calcium chloride melt, on the surface of titanium. Okabe, Devra, Oishi, Ono and Sadoway (Journal of Alloys and Compounds 237 (1996) 150) have deoxidised yttrium using a similar approach.

Ward *et al*, Journal of the Institute of Metals (1961) 90:6-12, describes an electrolytic treatment for the removal of various contaminating elements from molten copper during a refining process. The molten copper is treated in a cell with barium chloride as the electrolyte. The experiments show that sulphur can be removed using this process. However, the removal of oxygen is less certain, and the authors state that spontaneous non-electrolytic oxygen loss occurs, which may mask the extent of oxygen removal by this process. Furthermore, the process requires the metal to be molten, which adds to the overall cost of the refining process. The process is therefore unsuitable for a metal such as titanium which melts at 1660°C, and which has a highly reactive melt.

#### 25 Summary of Invention

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According to the present invention, a method for removing a substance (X) from a solid metal or semi-metal compound (M<sup>1</sup>X) by electrolysis in a melt of M<sup>2</sup>Y, comprises conducting the electrolysis under conditions such that reaction of X rather than M<sup>2</sup> deposition occurs at an electrode surface, and that X dissolves in the electrolyte M<sup>2</sup>Y.

According to one embodiment of the invention, M<sup>1</sup>X is a conductor and is used as the cathode. Alternatively, M<sup>1</sup>X may be an insulator in contact with a conductor.

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In a separate embodiment, the electrolysis product  $(M^2X)$  is more stable than  $M^1X$ .

In a preferred embodiment, M<sup>2</sup> may be any of Ca, Ba, Li, Cs or Sr and Y is Cl.

Preferably, M<sup>1</sup>X is a surface coating on a body of M<sup>1</sup>.

In a separate preferred embodiment, X is dissolved within M<sup>1</sup>.

In a further preferred embodiment, X is any of O, S, C or N.

In a still further preferred embodiment, M¹ is any of Ti, Si, Ge, Zr, Hf, Sm, U, Al, Ma, Nd, Mo, Cr, Nb, or any alloy thereof.

In the method of the invention, electrolysis preferably occurs with a potential below the decomposition potential of the electrolyte. A further metal compound or semi-metal compound (M<sup>N</sup>X) may be present, and the electrolysis product may be an alloy of the metallic elements.

The present invention is based on the realisation that an electrochemical process can be used to ionise the oxygen contained in a solid metal so that the oxygen dissolves in the electrolyte.

When a suitably negative potential is applied in an electrochemical cell with the oxygen-containing metal as cathode, the following reaction occurs:

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$$O + 2e^{-} \rightarrow O^{2}$$

The ionised oxygen is then able to dissolve in the electrolyte.

The invention may be used either to extract dissolved oxygen from a metal, i.e. to remove the a case, or may be used to remove the oxygen from a metal oxide. If a mixture of oxides is used, the cathodic reduction of the oxides will cause an alloy to form.

The process for carrying out the invention is more direct and cheaper than the more usual reduction and refining process used currently.

In principle, other cathodic reactions involving the reduction and dissolution of other metalloids, carbon, nitrogen, phosphorus, arsenic, antimony etc. could also take place. Various electrode potentials, relative to  $E_{Na} = O V$ , at  $700^{\circ}C$  in fused chloride melts containing calcium chloride, are as follows:

$$Ba^{2} + 2e^{-} = Ba \qquad -0.314 \text{ V}$$

$$Ca^{2} + 2e^{-} = Ca \qquad -0.06 \text{ V}$$

$$Hf^{4+} + 4e^{-} = Hf \qquad 1.092 \text{ V}$$

$$Zr^{4+} + 4e^{-} = Zr \qquad 1.516 \text{ V}$$

$$Ti^{4+} + 4e^{-} = Ti \qquad 2.039 \text{ V}$$

$$Cu^{4} + e^{-} = Cu \qquad 2.339 \text{ V}$$

$$Cu^{2} + 2e^{-} = Cu \qquad 2.92 \text{ V}$$

$$Cu^{2} + 4e^{-} = 20^{2} \qquad 2.77 \text{ V}$$

The metal, metal compound or semi-metal compound can be in the form of single crystals or slabs, sheets, wires, tubes, etc., commonly known as semi-finished or mill-products, during or after production; or alternatively in the form of an artefact made from a mill-product such as by forging, machining, welding, or a combination of these, during or after service. The element or its alloy can also be in the form of shavings, swarf, grindings or some other by-product of a fabrication process. In addition, the metal oxide may also be applied to a metal substrate prior to treatment, e.g. TiO<sub>2</sub> may be applied to steel and subsequently reduced to the titanium metal.

#### Description of the Drawings

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Figure 1 is a schematic illustration of the apparatus used in the present invention:

Figure 2 illustrates the hardness profiles of a surface sample of titanium before and after electrolysis at 3.0 V and 850°C; and

Figure 3 illustrates the difference in currents for electrolytic reduction of TiO<sub>2</sub> pellets under different conditions.

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#### Description of the Invention

In the present invention, it is important that the potential of the cathode is maintained and controlled potentiostatically so that only oxygen ionisation occurs and not the more usual deposition of the cations in the fused salt.

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The extent to which the reaction occurs depends upon the diffusion of the oxygen in the surface of the metal cathode. If the rate of diffusion is low, the reaction soon becomes polarised and, in order for the current to keep flowing, the potential becomes more cathodic and the next competing cathodic reaction will occur, i.e. the deposition of the cation from the fused salt electrolyte. However, if the process is allowed to take place at elevated temperatures, the diffusion and ionisation of the oxygen dissolved in the cathode will be sufficient to satisfy the applied currents, and oxygen will be removed from the cathode. This will continue until the potential becomes more cathodic, due to the lower level of dissolved oxygen in the metal, until the potential equates to the discharged potential for the cation from the electrolyte.

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This invention may also be used to remove dissolved oxygen or other dissolved elements, e.g. sulphur, nitrogen and carbon from other metals or semimetals, e.g. germanium, silicon, hafnium and zirconium. The invention can also be used to electrolytically decompose oxides of elements such as titanium, uranium, magnesium, aluminium, zirconium, hafnium, niobium, molybdenum, neodymium, samanium and other rare earths. When mixtures of oxides are reduced, an alloy of the reduced metals will form.

The metal oxide compound should show at least some initial metallic conductivity or be in contact with a conductor.

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An embodiment of the invention will now be described with reference to the drawing, where Figure 1 shows a piece of titanium made in a cell consisting of an inert anode immersed in a molten salt. The titanium may be in the form of a rod, sheet or other artefact. If the titanium is in the form of swarf or particulate matter, it may be held in a mesh basket. On the application of a voltage via a power source, a current will not start to flow until balancing reactions occur at both the anode and cathode. At the cathode, there are two possible reactions, the discharge of the cation from the salt or the ionisation and dissolution of oxygen. The latter reaction occurs at a more positive potential than the discharge of the metal cation and, therefore, will occur first. However, for the reaction to proceed, it is necessary for

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the oxygen to diffuse to the surface of the titanium and, depending on the temperature, this can be a slow process. For best results it is, therefore, important that the reaction is carried out at a suitably elevated temperature, and that the cathodic potential is controlled, to prevent the potential from rising and the metal cations in the electrolyte being discharged as a competing reaction to the ionisation and dissolution of oxygen into the electrolyte. This can be ensured by measuring the potential of the titanium relative to a reference electrode, and prevented by potentiostatic control so that the potential never becomes sufficiently cathodic to discharge the metal ions from the fused salt.

The electrolyte must consist of salts which are preferably more stable than the equivalent salts of the metal which is being refined and, ideally, the salt should be as stable as possible to remove the oxygen to as low as concentration as possible. The choice includes the chloride salts of barium, calcium, cesium, lithium, strontium and yttrium. The melting and boiling points of these chlorides are given below:

		Melting Point (°C)	Boiling Point (°C)
	BaCl <sub>2</sub>	963	1560
20	CaCl <sub>2</sub>	782	>1600
	CsCl	645	1280
	LiCl	605	1360
	SrCl <sub>2</sub>	875	1250
	YCl <sub>3</sub>	· <b>721</b>	1507

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Using salts with a low melting point, it is possible to use mixtures of these salts if a fused salt melting at a lower temperature is required, e.g. by utilising a eutectic or near-eutectic mixture. It is also advantageous to have, as an electrolyte, a salt with as wide a difference between the melting and boiling points, since this gives a wide operating temperature without excessive vaporisation. Furthermore, the higher the temperature of operation, the greater will be the diffusion of the oxygen in the surface layer and therefore the time for deoxidation to take place will be correspondingly less. Any salt could be used provided the

oxide of the cation in the salt is more stable than the oxide of the metal to be purified.

The following Examples illustrate the invention. In particular, Examples 1 and 2 relate to removal of oxygen from an oxide.

#### 5 Example 1

A white TiO<sub>2</sub> pellet, 5mm in diameter and 1mm in thickness, was placed in a titanium crucible filled with molten calcium chloride at 950°C. A potential of 3V was applied between a graphite anode and the titanium crucible. After 5h, the salt was allowed to solidify and then dissolved in water to reveal a black/metallic pellet. Analysis of the pellet showed that it was 99.8% titanium.

#### Example 2

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A strip of titanium foil was heavily oxidised in air to give a thick coating of oxide (c.50mm). The foil was placed in molten calcium chloride at 950°C and a potential of 1.75V applied for 1.5h. On removing the titanium foil from the melt, the oxide layer had been completely reduced to metal.

Examples 3 - 5 relate to removal of dissolved oxygen contained within a metal.

#### Example 3

Commercial purity (CP) titanium sheets (oxygen 1350-1450 ppm, Vickers Hardness Number 180) were made the cathode in a molten calcium chloride melt, with a carbon anode. The following potentials were applied for 3h at 950°C followed by 1.5h at 800°C. The results were as follows:

	V (volt)	Vickers	Oxygen
25		Hardness	Content
		Number	
	3 V	133.5	<200 ppm
	3.3 V	103	<200 ppm
	2.8 V	111	<200 ppm
30	3.1 V	101	<200 ppm

The 200 ppm was the lowest detection limit of the analytical equipment. The hardness of titanium is directly related to the oxygen content, and so measuring the hardness provides a good indication of oxygen content.

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The decomposition potential of pure calcium chloride at these temperatures is 3.2 V. When polarisation losses and resistive losses are considered, a cell potential of around 3.5V is required to deposit calcium. Since it is not possible for calcium to be deposited below this potential, these results prove that the cathodic reaction is:

$$O + 2e^{-} = O^{2-}$$

This further demonstrates that oxygen can be removed from titanium by this technique.

#### Example 4 10

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A sheet of commercial purity titanium was heated for 15 hours in air at 700°C in order to form an alpha case on the surface of the titanium.

After making the sample the cathode in a CaCl, melt with a carbon anode at 850°C, applying a potential of 3V for 4 hours at 850°C, the alpha case was removed as shown by the hardness curve (Figure 2), where VHN represents the Vicker's Hardness Number.

#### Example 5

A titanium 6 Al 4V alloy sheet containing 1800 ppm oxygen was made the cathode in a CaCl<sub>2</sub> melt at 950°C and a cathodic potential of 3V applied. After 3 hours, the oxygen content was decreased from 1800 ppm to 1250 ppm.

Examples 6 and 7 show the removal of the alpha case from an alloy foil. Example 6

A Ti-6A1-4V alloy foil sample with an alpha case (thickness about 40 µm) under the surface was electrically connected at one end to a cathodic current collector (a Kanthal wire) and then inserted into a CaCl, melt. The melt was contained in a titanium crucible which was placed in a sealed Inconel reactor that was continuously flushed with argon gas at 950°C. The sample size was 1.2 mm thick, 8.0 mm wide and ~50 mm long. Electrolysis was carried out in a manner of controlled voltage, 3.0V. It was repeated with two different experimental times and end temperatures. In the first case, the electrolysis lasted for one hour and the sample was immediately taken out of the reactor. In the second case, after 3 hours of electrolysis, the temperature of the furnace was allowed to cool naturally while maintaining the electrolysis. When the furnace temperature dropped to slightly lower than 800°C, the electrolysis was terminated and the electrode removed.

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Washing in water revealed that the 1 hour sample had a metallic surface but with patches of brown colour, whilst the 3 hour sample was completely metallic.

Both samples were then sectioned and mounted in a bakelite stub and a normal grinding and polishing procedure was carried out. The cross section of the samples was investigated by microhardness test, scanning electron microscopy (SEM) and energy dispersive X-ray analysis (EDX). The hardness test showed that the alpha case of both samples disappeared, although the 3 hour sample showed a hardness near the surface much lower than that at the centre of the sample. In addition, SEM and EDX detected insignificant changes in the structure and elemental composition (except for oxygen) in the deoxygenated samples.

#### Example 7

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In a separate experiment, Ti-6A1-4V foil samples as described above (1.2 mm thick, 8 mm wide and 25 mm long) were placed at the bottom of the titanium crucible which functioned as the cathodic current collector. The electrolysis was then carried out under the same conditions as mentioned in Example 6 for the 3-hour sample except that the electrolysis lasted for 4 hours at 950°C. Again using microhardness test, SEM and EDX revealed the successful removal of the alpha case in all the three samples without altering the structure and elemental composition except for oxygen.

Example 8 shows a slip-cast technique for the fabrication of the oxide electrode.

#### Example 8

A  $TiO_2$  powder (anatase, Aldrich, 99.9+% purity; the powder possibly contains a surfactant) was mixed with water to produce a slurry ( $TiO_2$ : $H_2O = 5:2$  wt) that was then slip-cast into a variety of shapes (round pellets, rectangular blocks, cylinders, etc) and sizes (from millimetres to centimetres), dried in room/ambient atmosphere ovemight and sintered in air, typically for two hours at 950°C in air. The resultant  $TiO_2$  solid has a workable strength and a porosity of 40~50%. There was notable but insignificant shrinkage between the sintered and unsintered  $TiO_2$  pellets.

0.3g-10g of the pellets were placed at the bottom of a titanium crucible containing a fresh CaCl<sub>2</sub> melt (typically 140g). Electrolysis was carried out at 3.0V (between the titanium crucible and a graphite rod anode) and 950°C under an argon environment for 5~15 hours. It was observed that the current flow at the

beginning of the electrolysis increased nearly proportionally with the amount of the pellets and followed roughly a pattern of 1 g TiO<sub>2</sub> corresponding to 1A initial current flow.

It was observed that the degree of reduction of the pellets can be estimated by the colour in the centre of the pellet. A more reduced or metallised pellet is grey in colour throughout, but a lesser reduced pellet is dark grey or black in the centre. The degree of reduction of the pellets can also be judged by placing them in distilled water for a few hours to overnight. The partially reduced pellets automatically break into fine black powders while the metallised pellets remain in the original shape. It was also noticed that even for the metallised pellets, the oxygen content can be estimated by the resistance to pressure applied at room temperature. The pellets became a grey powder under the pressure if there was a high level of oxygen, but a metallic sheet if the oxygen levels were low.

SEM and EDX investigation of the pellets revealed considerable difference in both composition and structure between metallised and partially reduced pellets. In the metallised case, the typical structure of dendritic particles was always seen, and no or little oxygen was detected by EDX. However, the partially reduced pellets were characterised by crystallites having a composition of Ca<sub>x</sub>Ti<sub>y</sub>O<sub>z</sub> as revealed by EDX.

#### 20 Example 9

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It is highly desirable that the electrolytic extraction be performed on a large scale and the product removed conveniently from the molten salt at the end of the electrolysis. This may be achieved for example by placing the TiO<sub>2</sub> pellets in a basket-type electrode.

The basket was fabricated by drilling many holes (~3.5 mm diameter) into a thin titanium foil (~1.0 mm thickness) which was then bent at the edge to form a shallow cuboid basket with an internal volume of 15x45x45 mm<sup>3</sup>. The basket was connected to a power supply by a Kanthal wire.

A large graphite crucible (140 mm depth, 70 mm diameter and 10 mm wall thickness) was used to contain the CaCl<sub>2</sub> melt. It was also connected to the power supply and functioned as the anode. Approximately 10g slip-cast TiO<sub>2</sub> pellets/blobs (each was about 10 mm diameter and 3 mm maximum thickness) were placed in the titanium basket and lowered into the melt. Electrolysis was conducted at 3.0V, 950°C, for approximately 10 hours before the furnace

temperature was allowed to drop naturally. When the temperature reached about 800°C, the electrolysis was terminated. The basket was then raised from the melt and kept in a water-cooled upper part of the Inconel tube reactor until the furnace temperature dropped to below 200°C before being taken out for analysis.

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After acidic leaching (HCl, pH<2) and washing in water, the electrolysed pellets exhibited the same SEM and EDX features as observed above. Some of the pellets were ground into a powder and analysed by thermo-gravitmetry and vacuum fusion elemental analysis. The results showed that the powder contained about 20,000 ppm oxygen.

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SEM and EDX analysis showed that, apart from the typical dendritic structure, some crystallites of  $CaTiO_x$  (x<3) were observed in the powder which may be responsible for a significant fraction of the oxygen contained in the product. If this is the case, it is expected that upon melting the powder, purer titanium metal ingot can be produced.

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An alternative to the basket-type electrode is the use of a "lolly" type TiO<sub>2</sub> electrode. This is composed of a central current collector and on top of the collector a reasonably thick layer of porous TiO<sub>2</sub>. In addition to a reduced surface area of the current collector, other advantages of using a lolly-type TiO<sub>2</sub> electrode include: firstly, that it can be removed from the reactor immediately after electrolysis, saving both processing time and CaCl<sub>2</sub>; secondly, and more importantly, the potential and current distribution and therefore current efficiency can be improved greatly.

#### Example 10

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A slurry of Aldrich anatase TiO<sub>2</sub> powder was slip cast into a slightly tapered cylindrical lolly (~20 nm length and ~ mm diameter) comprising a titanium metal foil (0.6 mm thickness, 3 mm width and ~40 mm length) in the centre. After sintering at 950°C, the lolly was connected electrically at the end of the titanium foil to a power supply by a Kanthal wire. Electrolysis was carried out at 3.0V and 950°C for about 10 hours. The electrode was removed from the melt at about 800°C, washed and leached by weak HCl acid (pH 1-2). The product was then analysed by SEM and EDX. Again, a typical dendritic structure was observed and no oxygen, chlorine and calcium could be detected by EDX.

The slip-cast method may be used to fabricate large rectangular or cylindrical blocks of TiO<sub>2</sub> that can then be machined to an electrode with a desired

shape and size suitable for industrial process. In addition, large reticulated TiO<sub>2</sub> blocks, e.g. TiO<sub>2</sub> foams with a thick skeleton, can also be made by slip cast, and this will help the draining of the molton salt.

The fact that there is little oxygen in a dried fresh CaCl<sub>2</sub> melt suggests that the discharge of the chloride anions must be the dominant anodic reaction at the initial stage of electrolysis. This anodic reaction will continue until oxygen anions from the cathode transport to the anode. The reactions can be summarised as follows:

10 anode:

cathode:

$$TiO_2 + 4e = Ti + 20^{2}$$

total:

$$TiO_2 + 4Cl' = Ti + 2Cl_2 + 2O^{2-}$$

When sufficient O2- ions are present the anodic reaction becomes:

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$$O^{2} = \frac{1}{2} O_2 + 2e^{-1}$$

and the overall reaction:

$$TiO_2 = Ti + O_2 t$$

Apparently the depletion of chloride anions is irreversible and consequently the cathodically formed oxygen anions will stay in the melt to balance the charge, leading to an increase of the oxygen concentration in the melt. Since the oxygen level in the titanium cathode is in a chemical equilibrium or quasi-equilibrium with the oxygen level in the melt for example via the following reaction:

It is expected that the final oxygen level in the electrolytically extracted titanium cannot be very low if the electrolysis proceeds in the same melt with controlling the voltage only.

This problem can be solved by (1) controlling the initial rate of the cathodic oxygen discharge and (2) reducing the oxygen concentration of the melt. The

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former can be achieved by controlling the current flow at the initial stage of the electrolysis, for example gradually increasing the applied cell voltage to the desired value so that the current flow will not go beyond a limit. This method may be termed "double-controlled electrolysis". The latter solution to the problem may be achieved by performing the electrolysis in a high oxygen level melt first, which reduces TiO<sub>2</sub> to the metal with a high oxygen content, and then transferring the metal electrode to a low oxygen melt for further electrolysis. The electrolysis in the low oxygen melt can be considered as an electrolytic refining process and may be termed "double-melt electrolysis".

Example 11 illustrates the use of the "double-melt electrolysis" principle. Example 11

A TiO<sub>2</sub> lolly electrode was prepared as described in Example 10. A first electrolysis step was carried out at 3.0V, 950°C overnight (~12 hours) in re-melted CaCl<sub>2</sub> contained within an alumina crucible.

A graphite rod was used as the anode. The lolly electrode was then transferred immediately to a fresh CaCl<sub>2</sub> melt contained within a titanium crucible. A second electrolysis was then carried out for about 8 hours at the same voltage and temperature as the first electrolysis, again with a graphite rod as the anode. The lolly electrode was removed from the reactor at about 800°C, washed, acid leached and washed again in distilled water with the aid of an ultrasonic bath. Again both SEM and EDX confirmed the success in extraction.

Thermo-weight analysis was applied to determine the purity of the extracted titanium based on the principle of re-oxidation. About 50 mg of the sample from the lolly electrode was placed in a small alumina crucible with a lid and heated in air to 950°C for about 1 hour. The crucible containing the sample was weighted before and after the heating and the weight increase was observed. The weight increase was then compared with the theoretical increase when pure titanium is oxidised to titanium dioxide. The result showed that the sample contained 99.7+% of titanium, implying less than 3000 ppm oxygen.

#### Example 12 30

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The principle of this invention can be applied not only to titanium but also other metals and their alloys. A mixture of TiO2 and Al2O3 powders (5:1 wt) was slightly moistened and pressed into pellets (20 mm diameter and 2 mm thickness) which were later sintered in air at 950°C for 2 hours. The sintered pellets were

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white and slightly smaller than before sintering. Two of the pellets were electrolysed in the same way as described in Example 1 and Example 3. SEM and EDX analysis revealed that after electrolysis the pellets changed to the Ti-Al metal alloy although the elemental distribution in the pellet was not uniform: the Al concentration was higher in the central part of the pellet than near the surface, varying from 12 wt% to 1 wt%. The microstructure of the Ti-Al alloy pellet was similar to that of the pure Ti pellet.

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Figure 3 shows the comparison of currents for the electrolytic reduction of  ${\rm TiO_2}$  pellets under different conditions. It can be shown that the amount of current flowing is directly proportional to the amount of oxide in the reactor. More importantly, it also shows that the current decreases with time and therefore it is probably the oxygen in the dioxide that is ionising and not the deposition of calcium. If calcium was being deposited, the current should remain constant with time.

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#### CLAIMS

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- 1. A method for removing a substance (X) from a solid metal, a metal compound or semi-metal compound (M<sup>1</sup>X) by electrolysis in a fused salt of M<sup>2</sup>Y or a mixture of salts, which comprises conducting the electrolysis under conditions such that reaction of X rather than M<sup>2</sup> deposition occurs at an electrode surface, and that X dissolves in the electrolyte M<sup>2</sup>Y.
- A method according to claim 1, wherein M<sup>1</sup>X is a conductor and is used as the cathode.
- A method according to claim 1, wherein M<sup>1</sup>X is an insulator and is used in 3. 10 contact with a conductor.
  - A method according to any preceding claim, wherein electrolysis is carried 4. out at a temperature of 700°C - 1000°C.
  - A method according to any preceding claim, wherein the electrolysis product (M<sup>2</sup>X) is more stable than M<sup>1</sup>X.
- A method according to any preceding claim, wherein M<sup>2</sup> is Ca, Ba, Li, Cs or 15 Sr and Y is Cl.
  - 7. A method according to any preceding claim, wherein M<sup>1</sup>X is a surface coating on a body of M1.
  - A method according to any of claims 1 to 6, wherein X is dissolved within 8. M¹.
  - A method according to any preceding claim, wherein X is O, S, C or N. 9.
  - A method according to any preceding claim, wherein M<sup>1</sup> is Ti or an alloy 10. thereof.
  - 11. A method according to any of claims 1 to 9, wherein M<sup>1</sup> is Si or an alloy thereof.
    - A method according to any of claims 1 to 9, wherein M<sup>1</sup> is Ge or an alloy 12. thereof.
    - 13. A method according to any of claims 1 to 9, wherein M<sup>1</sup> is Zr or an alloy thereof.
- A method according to any of claims 1 to 9, wherein M<sup>1</sup> is Hf or an alloy 14. 30 thereof.
  - A method according to any of claims 1 to 9, wherein M<sup>1</sup> is Sm or an alloy **15**. thereof.

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- 16. A method according to any of claims 1 to 9, wherein  $M^1$  is U or an alloy thereof.
- 17. A method according to any of claims 1 to 9, wherein M¹ is Al or an alloy thereof.
- 5 18. A method according to any of claims 1 to 9, wherein M¹ is Mg or an alloy thereof.
  - 19. A method according to any of claims 1 to 9, wherein M¹ is Nd or an alloy thereof.
  - 20. A method according to any of claims 1 to 9, wherein M<sup>1</sup> is Mo or an alloy thereof.
    - 21. A method according to any of claims 1 to 9, wherein M¹ is Cr or an alloy thereof.
    - 22. A method according to any of claims 1 to 9, wherein M¹ is Nb or an alloy thereof.
- 23. A method according to any preceding claim, wherein M¹X is in the form of a porous pellet or powder.
  - 24. A method according to any preceding claim, wherein electrolysis occurs with a potential below the decomposition potential of the electrolyte.
- 25. A method according to any preceding claim, wherein a further metal compound or semi-metal compound (M<sup>N</sup>X) is present, and the electrolysis product is an alloy of the metallic elements.

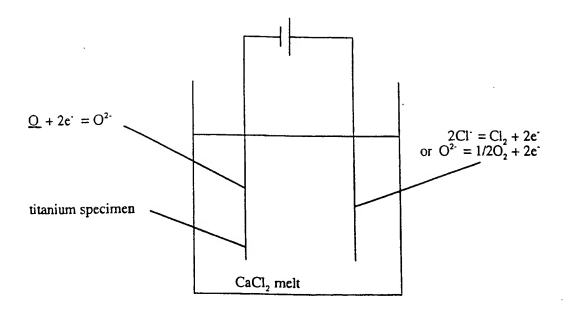


Figure 1

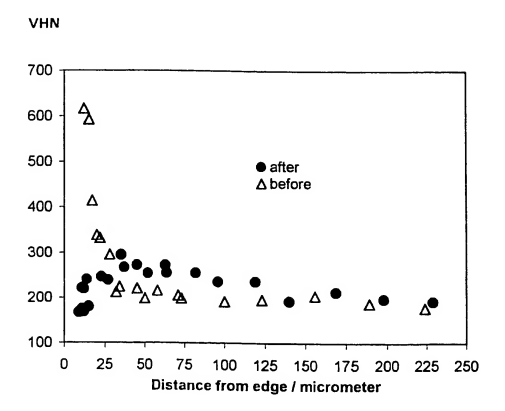


Figure 2

# Current/A

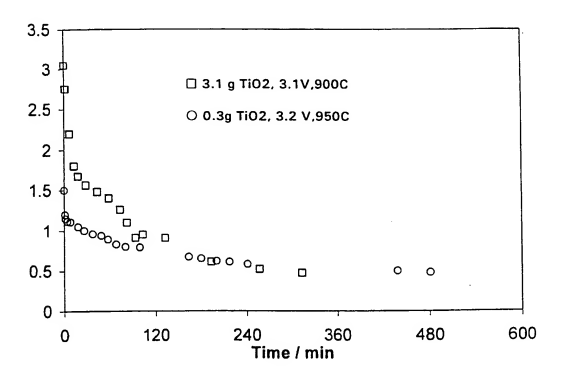


Figure 3

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